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Description

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This invention relates to a specific thermoplastic polymer composition which comprises an ethylene-propylene rubor, an ethylene copolymer, a propylene polymer (including a propylene-ethylene block copolymen) and tallsc, sowns good processability on injection modding, can give a molded product having a good appearance, has excellent surface hardness, impact resistance and adhesion of ceating, and thus is suited for the production of injection-molded products such as automobile parts.

Herelocies, many attempts have been made to enhance the value of molded products of various rubbers such as an ehylene-proylene copolymer rubber by improving heir fluidity and imparting rigidity to them. For instance, compositions prepared by incorporating polypropylene to rubber have been known as disclosed in Japanese Patent Publications Nos. 57-57049, 62-5460 and 62-5461. However, the proportion of polyprophene in these compositions is generally small, and polyprophien having high fluidity and high crystallinity is not particularly employed, so that the compositions have low crystallization rates. A long cooling time is therefore required when large-sized molded products of the compositions are produced by means of injection molding. The productivity is thus extremely low. In addition, since the compositions contain neither enthylene copolymers nor tale, they can give only such molded products that are poor in the surface smoothness and surface hardness.

The composition disclosed in Japanese Patent Publication No. 611-9651 comprises a relatively large amount of polypropylene. However, this composition also contains neither ethylene copplymers nor tate. Moreover, a patiely crosslinked rubber is used as a rubber component. For these reasons, the composition is also confronted with the same problems as in the above compositions in the productivity upon producing molded products, and in the surface smoothness and surface hardness of molded products.

On the other hand, Japanese Patent Publication No. 60-3420 discloses a composition prepared by incorporating an ethylene-proyelene ubber and talc into a propyleneethylene block copolymer. This composition is excellent in adhesion of coating and low-temperature impact resistance. Further, Japanese Patent Laid-Open Publication No. 1-204946 discloses a composition comprising an ethylene-propylene rubber, a propyleneethylene block copolymer, an ethylene copolymer and talc. This composition can give molded products having an improved dimensional stability. These compositions are, however, still insufficient in the surface hardness and smoothness of molded products. The molded products will be easily flawed due to their low surface hardness and have a poor appearance due to their poor surface smoothness.

It is therefore an object of the present invention to solve the above problems in the prior art and provide a thermoplastic polymer composition which shows good processability on injection molding, can give an injection-molded product having a good appearance and a low density, and is excellent in the surface hardness, impact resistance and achesion of coating.

SUMMARY OF THE INVENTION

It has now been found that the above object can be achieved by blending an ethylene-propylene rubber, an ethylene copolymer, a propylene polymer and tale in a specific proportion.

Thus, the present invention provides a thermoplastic polymer composition comprising (i) 100 parts by weight of a thermoplastic polymer component which comprises an ethylene-propylene rubber, an ethylene-α-olefin copolymer, a thermoplastic polymer composition comprising:

- (i) 100 parts by weight of a thermoplastic polymer component which comprises an ethylene-propylene rubber, an ethylene-coletin copolymer having a melting point determined by a differential scanning calorimeter of 65 to 90°C and a density of 0.87 to 0.89 g/cm³ in which the α-coletin has 4 to 8 carbon atoms and a propylene polymer (including an ethylene-propylene block copolymer), and
- (ii) 0 to 5 parts by weight of talc,

the thermoplastic polymer component being composed of, according to fractionation using o-dichlorobenzene as a solvent, component (A) which is a component soluble in the solvent at 40°C, component (B) which is a component insoluble in the solvent at 40°C but soluble at 110°C, and component (C) which is a component insoluble in the solvent even at 110°C in such a proportion that the total amount of the components (A) and (B) is from 50 to 70 parts by weight, the weight ratio of the component (A) to the component (B) being from 0.5 to 1.5, and the amount of the component (C) is from 50 to 30 parts by weight.

The composition of the present invention shows good processability when subjected to injection molding, can give molded products which have a good appearance, and is excellent in the surface hardness, impact resistance and adhasion of coatino.

More specifically, the present invention provides, for instance, a composition having a melt flow rate (MFR) of 10

g/10 min or more, a density of 0.95 g/cm³ or lower, a flexural modulus of 6,000 kg/cm² or more, a Flockwell hardness of 50 or more, an Izod impact strength at -30°C of 5 kg-cm/cm or more, and a peeling strength of a coated film, which will be described later. is 700 g/cm or more.

Moreover, since the proportion of a high-crystalline component (C) and a relatively high-crystalline component (B) in the composition of the present invention is larger than that in the conventional thermoplastic polymer compositions, the composition of the invention can be solidified by cooling in a shorter time than before. The cooling time required in the process of injection molding can thus be shortened. This brings about a remarkable increase in the production seed of molded products.

The composition of the present invention, which has the above-described advantageous properties, is suitably utilized for injection-molded automobile parts, especially for large-sized parts which needs coating and require a good appearance and flaw-resistance, such as a bumper, an air dam spoiler and a fascia.

The ethylene-propylene rubber for use in the present invention should have an MFR (at 23°C) of 0.3 to 3 g/10 min, preferably 0.5 to 2 g/10 min, preferably 0.5 to 2 g/10 min, and contain a proper amount of cystalline segment in the molecule. In the present invention, an ethylene-propylene rubber which contains 12% to 30% by weight, preferably 15% to 25% by weight, of propylene, and has a melting point determined by a differential scanning calorimeter of 30° to 60°C, preferably 35° bost 55°C, is preferred from the viewpoint of surface hardness and adhesion of costing. The ethylene-propylene rubber any ba an EPDM containing ethylidene nordomene, discipopantadiene, 1,4-haxadiene or the like as a third component. The ethylene-provylene rubber may also be a mixture of EPM and EPDM.

It is desirable that the ethylene-oc-olelin copolymer to be used in the present invention has an MFR (at 230°C) of 1 to 10 g/10 min, profilerably 2 to 8 g/10 min, from the viewpoint of processability and impact resistance. The ethylene-olelin copolymer has a melting point determined by a differential scanning calorimeter of 55° to 90°C, because it is important that the crystallinity of the ethylene-copolymer be not too high when the compatibility between the ethylene-propylene rubber and the ethylene-copolymer is taken into consideration, and a density of 0.87-0.89 g/cm³, from the viewpoint of surface hardness, impact resistance and adhesion of coating.

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The above-described athylene copolymer can be prepared by copolymerizing ethylene and an c-otelin in the presence of an ionic polymerization catalyst auch as a Elegiar catalyst, a Phillips catalyst or a Kaminsky catalyst. Producion methods applicable to the above copolymerization may be a gas phase fluidized bed method, a solution method, a slurry method, and a high pressure ionic polymerization method in which polymerization is carried out under a pressure of 200 kg/cm² or more at a temperature of 150° Cor higher. As long as the metting point of the resulting copolymer falls within the above range, the c-otelin content in the copolymer is not particularly limited; however, it is generally 12-30% by weight, preferably 1-52% by weight. The α-otelin to be copolymerized with athylena should be a 1-otelin having 4 to 8 carbon atoms. Examples of the 1-otelin include butene-1, 3-methylbutene-1, pentene-1, 4-methylpentene-1, haxene-1, heptene-1 and octene-1. Such α-otelins may be used singly or as a mixture of two or more.

From the viewpoint of processability and impact resistance, it is preferred that the propylene polymer for use in the present invention be a resin having an MFR (at 230°C) of 30 to 150 g/10 min, preferably 50 to 100 g/10 min, and containing 1.5% to 5% by Swy begint, preferably 2% to 7% by swight, of ethylene. It is further preferred that the resin contain a crystalline propylene polymer moiety with a density of 0.907 g/cm³ or more, preferably 0.908 g/cm³ or more, when surface hardness is taken into consideration. Among such preferable resins, a propylene-ethylene block copol-were is especially preferred.

The above mentioned MFR of the propylene polymer may be controlled at the time of polymerization, or adjusted by an organic peroxide such as a diacyl peroxide or a dialkyl peroxide after the polymerization is completed.

The propylene polymer may also be a copolymer with other unsaturated monomer such as maleic anhydride, methacrylic acid or trimethoxyvinylsilana which is introduced thereto by graft or random copolymerization. In particular, the use of a mixture of the propylene polymer and a crystalline polypropylene grafted with maleic anhydride or trimethoxyvinylsilane can improve the surface hardness of the resulting composition.

A stereospecific catalyst is employed for the production of the above propylene polymer. Typical preparation methods of the catalyst are a method as disclosed in Japanese Plant Laid-Copen Publication Nos. 56: 100806. 56: 12020.

38-104907, in which a titanium trichloride composition prepared by reducing ittanium tetrachloride with an organoaluminum compound and then treating with various electron donors and electron acceptors is combined with an organoaluminum compound and an airomatic carboxylis acid ester, and a method as disclosed in Japanese Patent Laid-Open Publications Nos. 57-63310, 63-43915 and 63-83115, in which titanium tetrachloride and various electron donors are brought into contact with a magnesistum halide to give a carrier-type catalyst.

The above three kinds of polymer components are mixed so that the resulting polymer mixture (thermoplastic polymer component) can be composed of, according to fractionation using o-dichlorobenzene as a solvent, component (a) which is a component soluble in the solvent at 40°C, component (B) which is a component insoluble in the solvent at 40°C but soluble at 110°C, and component (C) which is a component insoluble in the solvent aven at 110°C in such a proportion that the total amount of the components (A) and (B) is from 50 to 70 parts by weight, the weight ratio of the component (A) to the component (B) is from 50 to 30 parts

by weight.

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In the case where the total amount of the components (A) and (B) is less than the above range, that is, the amount of the component (C) is in excess of the above range, the resulting composition has a poor impact resistance. On the other hand, when the total amount of the components (A) and (B) is more than the above range, the resulting composition cannot have a sufficiently high flourual modulus. Further, when the weight ratio of the component (A) to the component (B) (Component (A)component (B)) is less than the above range, the resulting composition will exhibit a poor adhesion of coating; while when the weight ratio is in excess of the above range, a molded product of the composition will exhibit a

When the intrinsic viscosity ([η]) of the component (C) is high, the resulting composition requires a high molding temperature, leading to an increase in molding yellow. It is therefore desirable that the intrinsic viscosity of the component (C) be 2.0 d/g or less, preferably 1.7 d/g or less.

The crystallinity of the propylene polymer can be shown by the proportion of highly-crystalline polypropylene, an index of which can be given by the following formula:

Amount of component (C) (Amount of component (C) + Amount of component (D))

wherein component (D) indicates a polypropylene moiety contained in the component (B), determined by ¹³C-NMR. In the present invention, it is preferred that the above index be 0.70 or more, preferably 0.75 or more, from the viewpoint of surface hardness

In the present invention, use may be made of a mixture of two or more kinds of the ethylene-propylene rubber, two more kinds of the ethylene copyloner, and two or more kinds of the propylene polymor, as long as the components (A), (B), (C) and (D) of the resulting thermoplastic polymer component can satisfy the allorementioned conditions.

The taic usable in the present invention should preferably have an average particle size of 5.0 µm or less, preferably from 0.5 to 3.0 µm, and a specific surface area of 3.5 m³/g or more, preferably from 3.5 to 6.0 m³/g. Such taic may be prepared by a dry pulverization and the subsequent dry classification. When the average particle size of talc is in excess of 5.0 µm, the resulting composition is fikely to exhibit a poor impact resistance. The average particle size of talc herein is a particle size at a cumulative amount of 50% by weight in a cumulative particle size distribution curve obtained by a liquid phase sedimentation light transmission method using, for instance, Model CP manufactured by Shimaddu Corp. The specific surface area can be measured by an air permeation method using, for instance, a specific surface area measuring apparatus Model SP from amufactured by Shimaddu Corp.

To impart high adhesion of ceating to a molded product of the composition of the invention, it is preferable to control the amount of a non-crystalline moiety in the molded product to 50% by weight or more of the total amount of the resin components contained therein. Specifically, it is preferable that the thermoplastic polymer component comprising the athylene-propylene rubber, the athylene copolymer and the propylene polymer have a degree of crystallinity, determined by oulsed MMR. of 50% or less.

The amount of talc to be used is from 0 to 5 parts by weight, for 100 parts by weight of the total amount of the ethylene-propylene rubber, the ethylene copolymer and the propylene polymer.

The talc can be used without subjecting to any treatment. However, in order to improve the adhesion between the talc and the polymers, or the dispersibility of the talc in the polymers, the talc may be treated with various organic titanate coupling agents, silane coupling agents, filant parties, talty acids, metal salts of fatty acid, fatty acid elsters, and the like.

Other auxiliary components may be added to the composition of the present invention unless they substantially impair the advantageous properties of the composition.

Examples of the auxiliary components usable in the present invention include additives which are conventionally employed in thermoplastic polymer compositions, for example, a processing stabilizer, an antioxidant, an ultraviolet absorbor, a light stabilizer, various scaps such as motal scaps, an antilast, a flubricant, a nucleator, a pigment and a dispersant for pigment. In addition, whiskers such as fibrous potassium titanate, fibrous magnesium oxalfate and fibrous aluminum borate, and carbon fibers, which are known as materials capable of imparting higher flexural modulus than that imparted by table, can be emplowed, if necessing.

The composition of the present invention can be prepared by kneading the polymer components, the talc, and, if necessary, the auxiliary components by any of an ordinary extruder, a Banbury mixer, a roller, a Brabender and a kneader. However, the use of a twin-screw extruder is desirable in the present invention.

From the composition of the present invention, molded products may be prepared by any known molding method set as an injection molding method, no extrusion molding method and a blow molding method. However, an injection molding method will be most advantageously employed in view of the inherent properties of the composition.

This invention will now be explained in more detail with reference to the following examples, which are given merely for illustration of this invention and are not intended to be limiting thereof.

In the examples, solvent fractionation was carried out in the following manner:

- (1) 5 g of a sample and 1.5 g of 2,6-di-t-butyl-p-phenol as an antioxidant were dissolved in 1.5 ℓ of o-dichlorobenzane at 140°C. The resulting mixture was filtrated through a 0.45-µm Teffon filter at 140°C to remove insoluble components such as a filter.
- (2) After redissolving the filtrate at 140°C, 300 g of Celite (#545) was added to the solution. While stirring, the resulting mixture was cooled to room temperature at a cooling rate of 10°C/hour to provide a coating on the surface of the Celite.
 - (3) The coated Celite was filled in a cylindrical column. To this column, o-dichlorobenzene containing the above antioxidant in the same concentration as above was introduced, and fractionation was carried out by heating the column to temperatures of 40°C, 110°C and 140°C thereby to olute the coating.
 - (4) After the fractionation was completed, a large amount of methanol was added to each fraction, followed by filtration through a 0.45-µm Tellon filter. After drying in vacuum, each fraction was weighed. The proportion of each fraction was determined on the basis of the total weight of the three fractions.

Measuring methods used in the examples are as follows:

- (i) Melting Point: 10 mg of a sample was placed in a differential scanning calorimeter, for instance, Model 910 manufactured by Du Pont Corp. After heating to a temperature of +160°C, the sample was cooled to -100°C at a constant cooling rate of 10°C/min. Thereafter, the sample was heated again at a constant heating rate of 20°C/min. A temperature corresponding to the peak of the thermogram obtained was taken as the melting point of the sample.
- (ii) Intrinsic Viscosity: Ascosity: Associative and in ordinations of various concentrations, reagner containing 0.2% by well of the above-mentioned in admitted place of various concentrations, reagnering from 0.1 to 3.9th, of the polymer mer. Measurements of viscosity were carried out at a temperature of 1.40°C. The intrinsic viscosity of the polymer was determined by a viscosity were to a point of 2.50°C.
- (iii) Propylene Content in Component (B): The propylene content was determined by an integrated intensity of signals deriving from a polypropylene carbon, which appear in the vicinity of 46.5 ppm from TMS (tetramethylsillane) in a 19C-MMR spectrum.
 - (iv) Total Crystallinity of Resin: Determined by pulsed NMR (see Kobunshi Jikk engaku 18, "Magnetic Resonance of Polymer", pp. 143-144, Kyoritsu Shuppan Kabushiki Kaisha).
- 30 (v) MFR: Measured in accordance with ASTM-D1238 with application of a load of 2.16 kg at a temperature of 230°C.
 - (vi) Density: Measured in accordance with ASTM-D1505 at a temperature of 23°C.
 - (vii) Flexural Modulus: Measured in accordance with ASTM-D790 at a temperature of 23°C.
 - (viii) Surface Hardness: Evaluated by a Rockwell hardness (Scale R), measured in accordance with ASTM-D785 at a temperature of 23°C.
 - (ix) Impact Resistance: Evaluated by an Izod value at a temperature of -30°C, measured in accordance with ASTM-D256.
 - (x) Adhesion of Coating: Evaluated by the peeling strength of a coated film, determined in accordance with the following manner:

40 <1> Coating Method

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- a. An injection-molded specimen was exposed to the vapor of boiling 1,1,1-trichloroethane for 30 seconds, and then allowed to stand at room temperature for 30 minutes for drying.
- b. The lower half of the surface of the specimen was covered with a masking tape while the upper half thereof remained uncovered.
 - c. A polyurethane-modified polyolelin primer for polypropylene, "Soflex 2500" manufactured by Kansai Paint Co., Lid., was coaled not he specimen by means of spray coaling to form a primer layer with a thickness of approximately 10 µm. After drying the primer at room temperature for 30 minutes, the masking tape was peeled off the seecimen.
- d. A one-can urethane coating containing an isocyanate hardening agent, "Flexen 105" manufactured by Nihon B-Chemical K.K., was then coated onto the specimen by means of spray coating to form a coating layer with a thickness of approximately 60 µm. The specimen was placed in an air oven adjusted to a temperature of 120°C for 30 minutes to bake the coating, and then allowed to stand at room temperature for 48 hours. The specimen thus obtained was used in the peeling strength test described below.

<2> Measurement

a. A cellophane adhesive tape was adhered to the entire surface of the test specimen obtained above. On the

surface of the adhesive tape, cuts reaching to the base were made by a cutter in the long direction of the specimen at 10-mm intervals.

b. The coating layer formed on the surface not coated with primer was peeled, together with the cellophane adhesive tape adhered thereon, off the specimen, and bent to the 190-degree direction. The specimen in this state was set in a topic to state.

c. The tester was operated at 23°C at a rate of pulling of 50 mm/min. From the curve obtained on the recorder, a value corresponding to the peak was obtammed. The average of such values for ten peaks was taken herein as the value indicating the peeling strength of coating of the specimen.

10 Examples 1 and 2 and Comparative Examples 1 to 6

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Materials shown in Table 1 were mixed in accordance with the formulation shown in Table 2. To the resulting mixers, 0.1 part by weight of 2.6-dit-bulyl-2-hydroxyphenylpropionatelymethane and 0.5 part by weight of carbon black were added, and mixed by a "Super Mixer" manufactured by Kawada Seisakusho K.K. for 5 minutes. The mixtures thus obtained were kneaded and granulated at 210°C by a winn-screw kneaded. "FOM" manufactured by Kobe Steel L.U.1. Gold where more black were kneaded and granulated at 210°C by a winn-screw kneaded." FOM manufactured by Kobe Steel L.U.1. Gold where more slets to obview compositions.

Injection-moded specimens of the thus obtained thermoplastic polymer compositions were respectively prepared by an injection molding machine with a clamp pressure of 100 ton at a molding temperature of 220°C. The properties of the specimens were evaluated in accordance with the above-described measuring methods. Furthermore, evaluation of the appearance (surface smoothness) of molded products was made in the following manner:

Automobile bumpers (weight: 5 kg) were prepared by an injection molding machine with a clamp pressure of 4,000 ton at a molding temperature of 220°C. On the bumpers thus obtained, coating was conducted in the following manner:

- a. The bumper was exposed to the vapor of boiling 1,1,1-trichloroethane for 60 seconds.
- b. A chlorinated polypropylene primer for polypropylene, "R-117" manufactured by Nihon B-Chemical K.K., was then coated onto the bumper by means of spray coating to form a primer layer with a thickness of approximately 15 µm. The bumper was placed in an air oven edupated to a temperature of 80° C or 10 minutes to 8 was the coating.
 c. A melamine-acrylate coating, "R-320" manufactured by Nihon B-Chemical K.K., was then coated onto the bumper by means of spray coating to form a coating layer with a thickness of approximately 35 µm. The bumper was placed in an air oven at 120°C for 20 minutes to bake the coating.

The surface smoothness of the coated bumper was evaluated by visual observation in comparison with a sheet metal having the same coating.

The results are shown in Tables 3 and 4.

Table 1

		Table	1	
Material	s for Use in Examp	les and Comparative Examples		
<ethyler< th=""><th>ne-Propylene Rubb</th><th>91></th><th></th><th></th></ethyler<>	ne-Propylene Rubb	91>		
Туре	MFR (g/10 min)	Melting Point	Propylene Content (% by weight)	
EPR-1	1.0	79		12
EPR-2	1.8	53		16
EPR-3	0.6	39		24
EPR-4	0.9	19		32
<ethyler< td=""><td>ne Copolymer></td><td></td><td></td><td></td></ethyler<>	ne Copolymer>			
Туре	MFR (g/10 min)	Melting Point (°C) Density (g/cm		Butene Content (% by weight)
PEX-1	3.5	105	0.915	10
PEX-2	2.3	88	0.900	15
PEX-3	7.5	68	0.882	24
PEX-4	5.2	55	0.890	32

Table 1 (continued)

		Table 1 (contin	nued)	
Materia	ls for Use in Examp	oles and Comparative Examples		
<propyl< th=""><th>ene-Ethylene Block</th><th>Copolymer></th><th></th></propyl<>	ene-Ethylene Block	Copolymer>		
Туре	MFR (g/10 min) Ethylene Content (% by weight) Density of Propylene Polymer Moiety			
PP-1	16	4.5	0.907	
PP-2	55	7.2	0.905	
PP-3	53	4.0	0.908	
PP-4	87	5.5	0.907	
PP-5	142	2.7	0.911	
<talc></talc>				
Туре	Average Particle Size (μm)		Specific Surface Area (m²/g)	
Talc-1	2.1		4.1	
Talc-2	6.7		2.6	

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Table 2

Formulation								
	EPR		PEX		PP		Talc	
	Туре	Parts by Weight	Туре	Parts by Weight	Туре	Parts by Weight	Туре	Parts by Weight
Ex.1	EPR-3	15	PEX-3	15	PP-3	70	Talc-1	5
Ex.2	EPR-3	15	PEX-3	20	PP-3	65	Talc-1	5
Comp. Ex. 1	EPR-1	25	PEX-1	25	PP-3	50	Talc-1	5
Comp. Ex. 2	EPR-4	10	PEX-4	10	PP-3	80	Talc-1	5
Comp. Ex. 3	EPR-3	5	PEX-2	30	PP-3	65	Talc-1	5
Comp. Ex. 4	EPR-3	30	PEX-2	5	PP-5	65	Talc-1	5
Comp. Ex. 5	EPR-3	20	PEX-2	15	PP-2	65	Talc-1	10
Comp. Ex. 6	EPR-3	20	PEX-2	15	PP-1	65	Talc-2	5

Table 3

Examples		
	1	2
component (A) + component (B) parts by weight	54	62
component (A)/component (B)	0.8	0.8
[η] of component (C) dl/g	1.2	1.2
Proportion of high-crystalline PP (*1)	0.82	0.83
Total crystallinity of resin %	47	44
MFR g/10 min	21	24
Density g/cm ³	0.93	0.93
Flexural modulus Kg/cm ²	8,800	8,200
Rockwell hardness	64	60

^{*1:} Proportion of high-crystalline PP = Amount of component (C)/(Amount of component (C) + Amount of component (D))

Table 3 (continued)

Examples					
	1	2			
Izod value (at -30°C) Kg-cm/cm	5.6	8.5			
Peeling strength of coating g/cm	800	950			
Surface smoothness (*2)	0	0			
Surface smoothness (*2) (Note)	0				

^{*2: ...} Superior to sheet metal

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Comparative Examples							
	1	2	3	4	5	6	
component (A) + component (B) parts by weight	74	46	59	55	61	60	
component (A)/component (B)	1.2	0.9	0.3	2.3	1.4	1.2	
[η] of component (C) dl/g	1.3	1.1	1.5	0.8	1.6	2.3	
Proportion of high-crystalline PP (*1)	0.81	0.80	0.80	0.86	0.67	0.78	
Total crystallinity of resin %	37	47	43	53	41	45	
MFR g/10 min	16	26	27	34	19	6.9	
Density g/cm ³	0.93	0.93	0.93	0.93	0.96	0.93	
Flexural modulus Kg/cm²	5,000	10,600	8,600	7,500	9,300	7,30	
Rockwell hardness	51	48	65	52	45	53	
Izod value (at -30°C) Kg-cm/cm	8.2	4.2	6.0	5.3	6.6	4.8	
Peeling strength of coating g/cm	450	760	390	520	880	800	
Surface smoothness (*2)	0	0	0	×	0	0	

^{*1:} Proportion of high-crystalline PP = Amount of component (C)/(Amount of component (C) + Amount of component (D))

Cleims

1. A thermoplastic polymer composition comprising:

(i) 100 parts by weight of a thermoplastic polymer component which comprises an ethylene-propylene rubber, an ethylene-a-olelin copolymer having a melting point determined by a differential scanning calorimeter of 65 to 90°C and a density of 0.87 to 0.88 g/cm³ in which the α-olelin has 4 to 8 carbon atoms and a propylene polymer (including an ethylene-propylene block copolymer), and

(ii) 0 to 5 parts by weight of talc,

the thermoplastic polymer component being composed of, according to fractionation using o-dichlorobenzene as a solvent, component (A) which is a component soluble in the solvent at 40°C, component (B) which is a comcomponent insoluble in the solvent at 40°C but soluble at 110°C, and component (O) which is a component insoluble in the solvent even at 110°C in such a proportion that the total amount of the components (A) and (B) is from 50 to 70 parts by weight, the weight ratio of the component (A) to the component (B) being from 0.5 to 1.5, and the amount of the component (C) is form 50 to 30 parts by weight.

O ... Almost equal to sheet metal × ... Inferior to sheet metal

^{*2: @...} Superior to sheet metal

O ... Almost equal to sheet metal × ... Inferior to sheet metal

- The thermoplastic polymer composition according to claim 1, wherein the ethylene-propylene rubber has a melting
 point of 30°C to 60°C, when measured by a differential scanning calorimeter.
- The thermoplastic polymer composition according to claim 1 or 2, wherein the component (C) has an intrinsic viscosity of 2 dl/g or less.
- The thermoplastic polymer composition according to any one of the preceding claims, wherein the amount of the
 component (C), and that of component (D) which is a polypropylene moiety contained in the component (B), determined by "32-NMR, aslist withe following conditions:

Amount of the component (C)/ (Amount of the component (C) + Amount of the component (D)) > 0.7.

- The thermoplastic polymer composition according to any one of the preceding claims having a degree of crystallinity, determined by pulsed NMR, of 50% or less.
- The thermoplastic polymer composition according to any one of the preceding claims, wherein the talc has an
 average particle size of 5 μm or less and a specific surface area of 3.5 m²/g or more.
- The thermoplastic polymer composition according to any one of the preceding claims having a flexural modulus
 of 6,000 kg/cm² or more.

Revendications

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- 1. Composition polymère thermoplastique comprenant :
 - (i) 100 parties en poids d'un constituan polymère thermoplastique qui comprend un caoutchouc éthylène-propylène, un copolymère éthylène-az-olétine ayant un point de fusion déterminé par calorimétrie par analyse différentielle de 55 à 90°C, et une masse volumique de 0,07° à 0,88 g/cm², où l'ez-olétine a de 4 à 8 atomes de carbone, et un polymère du propylène (comprenant un copolymère séquencé éthylène-propylène), et (iil de 0 à 5 parties en poids de talsc.

le constituant polymère thermoplastique étant, selon un fractionnement utilisant de l'o-dichlorobarzène en tuque solvant, constitué du constituant (A), qui est un constituant soluble dans le solvant à 40°C, d'un constituant (B) qui est un constituant nonstituant insoluble dans le solvant à 40°C, mais qui y est soluble à 110°C, et d'un constituant (C), qui est un constituant insoluble dans le solvant même à 110°C, selon une proportion telle que la quantité totale des constituants (A) et (B) soit de 50 à 70 parties en poids, le rapport pondéral du constituant (A) au constituant (B) étant de 0,5 à 1,5, et que la quantité du constituant (C) soit de 50 à 30 parties en poids.

- Composition polymère thermoplastique selon la revendication 1, dans laquelle le caoutchouc éthylène-propylène a un point de fusion de 30 à 60°C, quand il est mesuré par calorimétrie par analyse différentielle.
- Composition polymère thermoplastique selon la revendication 1 ou 2, dans laquelle le constituant (C) a une viscosité intrinsèque de 2 dl/g ou moins.
 - Composition polymère lihermoplastique selon l'une quelconque des revendications précédentes, dans laquelle la quantité du constituant (C) et celle du constituant (D), quil est un fragment polypropylène contenu dans le constituant (B), satisfont, la détermination utilisant une RIMN-19C, aux conditions suivantes:

quantité du constituant (C)/ (quantité du constituant (C) + quantité du constituant (D)) > 0,7.

- Composition polymère thermoplastique selon l'une quelconque des revendications précédentes, ayant un degré de cristallinité, déterminé par RMN pulsée, de 50 % ou moins.
 - Composition polymère thermoplastique selon l'une quelconque des revendications précédentes, dans laquelle le talc a une granulométrie moyenne de 5 μm ou moins, et une aire spécifique de 3,5 m²/g ou plus.

 Composition polymère thermoplastique selon l'une quelconque des revendications précédentes, ayant un module d'élasticité en flexion de 6000 kg/cm² ou plus.

5 Patentansprüche

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- Thermoplastische Polymerzusammensetzung, umfassend:
 - (i) 100 Gew Teile einer thermoplestischen Polymerkomponente, die ein Ethylen-Propylen-Gummi, ein Ethylen-Olleflosopphierr mit einem Schmelzpunkt, bestimmt durch Differentialskeinreinierie, von 65 bis 90°C und einer Dichte von 0,87 bis 0,89 g/cm³, worin das A-Olefin 4 bis 8 Köhlenstoflatome aufweist, und ein Propylengummi (einschließlich einem Ethylen-Propylens-Dickockophyme umfaßt, und
 - (ii) 0 bis 5 Gew.Teile Talkum.

worin sich die Ihermoglastische Polymerkomponente entsprechend der Franktionierung unter Verwendung von o-Dichlorbanzol als Lösungsmittel aus der Komponente (A), die eine Komponente ist, die in dem Lösungsmittel bei 40°C löslich ist, einer Komponente (B), die eine Komponente ist, die in dem Lösungsmittel bei 40°C unlöslich, aber bei 110°C idslich ist, und einer Komponente (C), die eine Komponente ist, die in dem Lösungsmittel sellst bei 110°C unlöslich ist, in einem sochen Anteil zusammensetzt, daß die Gassamtnenge der Komponenten (A) von (B) von 50 bis 70 Gew. Failen, das Gewichtsverhältnis der Komponente (A) zu der Komponente (B) von 0,5 bis 1,5 und die Menge der Komponente (C) S0 bis 30 Gew Teile ist.

- 2. Thermoplastische Polymerzusammensetzung nach Anspruch 1, worin der Ethylen-Propylen-Gummi einen Schmelzpunkt von 30 bis 60°C hat, gemessen durch ein Differentialkalorimeter.
 - Thermoplastische Polymerzusammensetzung nach Anspruch 1 oder 2, worin die Komponente (C) eine Intrinsikviskosität von 2 dl/g oder weniger hat.
- Thermoplastische Polymerzusammensetzung nach einem der vorhergehenden Ansprüche, worin die Menge der Komponente (C) und die der Komponente (D), die ein Polymopylenanteil ist, der in der Komponente (B) enthalten ist, bestimmt durch ¹G-NMR, die folgenden Bedringungen erfüllt:

Menge der Komponente (C) / (Menge der Komponete (C) + Menge der Komponente (D)) > 0,7.

- Thermoplastische Polymerzusammensetzung nach einem der vorhergehenden Ansprüche, mit einem Kristallinitätsorad, bestimmt durch Puls-NMR, von 50% oder weniger.
- 49 6. Thermoplastische Polymerzusammenseitzung nach einem der vorhergehenden Ansprüche, worin das Talkum eine durchschnittliche Teilchengröße von 5 µm oder weniger und eine spezifische Oberfläche von 3,5 m²/g oder mehr hat.
 - Thermoplastische Polymerzusammensetzung nach einem der vorhergehenden Ansprüche mit einem Biegemodul von 6000 kg/cm² oder mehr.